Silylation **Chlorides** Palladium-catalyzed of Bulky Arvl with **Dialkoxydisilanes** Keitaro Fukui^a

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Abstract Arylsilanes bearing a bulky alkoxy group on the silicon were synthesized from aryl chlorides and dialkoxydisilanes under reaction conditions utilizing SingaCycle-A3 as a palladium precatalyst and lithium benzoate in wet DMA. This report proposes the first direct and catalytic method for introducing tert-butoxy- or 1-adamantyloxysilyl groups onto various aryl moieties through the silylation reaction.

Key words palladium, alkoxysilane, disilane, arylsilane,

Due to their unique chemical reactivity and potential synthetic utility, organosilicon compounds have frequently been the target of synthetic studies.1 In the current framework of organic synthesis, silicon functional groups should become more balanced in terms of stability and ease of activation. Some are rather too stable for facile transformations; others are too labile to manage during standard synthetic manipulations. Therefore, prompting better balances between stability and ease of activation would widen the possibility of using silyl groups as alternative key functional groups in organic synthesis. One of the promising solutions to this dilemma is the use of bulky alkoxy substituents to balance them.² Our research focus has thus turned to a tBuOMe₂Si^{3,4} group, as a candidate for such a balanced functional group. The special feature of this silyl group is its exceptional stability against water or bases in contrast to other labile primary or secondary alkoxy silyl groups. This silyl group is therefore expected to serve as a reliable functional group that can survive during multistep synthetic transformations. We accordingly focused on the development of a general method to introduce a *t*BuOMe₂Si group via silvlation of aryl halides. Typical silvlation reagents for aryl halides include hydrosilanes,5 silylboranes,6 and disilanes.5d,7 We chose to employ the corresponding disilane 5 as the silicon source for our crosscoupling-type silvlation reaction. Compound 5 can be easily prepared in one step^{3b} and this reagent is now commercially

available. Among the previously reported conditions for silylation between an aryl halide and a dialkoxydisilane5d,7g,7i-l, Denmark's seminal work was the first and only method to introduce an alkoxysilyl group (Scheme 1a) wherein aryl bromide 1 reacted with diethoxydisilane 2 using a PdCl₂/JohnPhos catalytic system to give arylsilane 3.5d As such, introduction of a silyl group with a more bulky tertiary alkoxy group such as tBuO has yet to be reported. Herein, we report our research on a palladium-catalyzed method for introducing a *t*BuOMe₂Si group through the silvlation of aryl chloride with the corresponding disilane 5 (Scheme 1b).



Scheme 1 Catalytic Alkoxysilylation of Aryl Halide with Dialkoxydisilane

As a result of our optimization of palladium-catalyzed silylation of ethyl *p*-chlorobenzoate 4a with disilane 5, we eventually devised the standard conditions for the reaction: 4a (0.17 M), 1.5 equiv disilane 5, 3 mol% SingaCycle-A38, 1.5 equiv LiOBz , 5 mol% H₂O, DMA, 100 °C, 12 h.9 Table 1 shows results associated with differences based on the deviations listed. Under the standard conditions, arylsilane **6a** was obtained in 75% NMR yield (75% isolated yield) along with the formation of the corresponding homo-coupling product, diethyl [1,1'-biphenyl]-4,4'-dicarboxylate, in 4% yield. Pd-PEPPSI-IPr¹⁰ was also applicable albeit with a slower reaction rate (entry 2). When the ligand was changed to *t*BuDavePhos or CyJohnPhos, the reaction became even more sluggish (entries 3, 4). With JohnPhos that gave a good result in Denmark's report,^{5d} a low conversion

Table 1 Optimization of Silylation Conditions					
EtO ₂ C Cl fBuO fDO				6a	∕ Si∼OfBu
Entry	Deviations from the standard conditions		Yield (%)ª		
			4a	6a	biaryl
1	None		0	75	4
				(75 [♭])	
2	Pd-PEPPSI-IPr instead of SingaCycle-A3		7	65	2
3	3.0 mol% tBuDavePhos, 1.5 mol% Pd ₂ (dba) ₃		20	48	3
	instead of SingaCy				
4	3.0 mol% CyJohnP	12	49	6	
F	2.0 mol% lobpPho	57	17	-1	
5	instead of SingaCv	cle-A3	57	17	<1
6	0.1 equiv instead o	59	13	0	
7	Et ₃ N instead of LiOBz		84	3	0
8	LiOAc instead of LiOBz		12	55	1
9	LiOPiv instead of LiOBz		5	53	1
10	Li ₂ CO ₃ instead of LiOBz		68	12	0
11	NaOBz instead of LiOBz		0	54	11
12	KOBz instead of Li	0	31	43	
13	NMP instead of DN	0	71	<1	
14	DMF instead of DN	29	50	3	
15	1,4-dioxane instea	90	0	0	



 $^o\text{Determined}$ by ^1H NMR analysis using 1,3,5-trimethoxybenzene or mesitylene as an internal standard. $^b\text{Isolated}$ yield.

Table 2 Scope of Reaction with Respect to Aryl Chloride^{a,b}



^aisolated yield. ^bConditions: **4** (0.5 mmol, 0.17 M), **5** (1.5 equiv), SingaCycle-A3 (3 mol%), LiOBz (1.5 equiv), H₂O (5 mol%), DMA, 100 °C, 12–60 h. ^cDetermined by ¹H NMR analysis using 1,3,5-trimethoxybenzene mesitylene as an internal standard. ^dSingaCycle-A3 (6 mol%).

was observed (entry 5). The amount of the base heavily affected the reaction outcome. With 0.1 equiv of LiOBz, the minimum necessary amount for activating SingaCycle-A3, only a 13% yield of the product was obtained (entry 6). This is in contrast with our previous report on the base-free silylation of aryl chloride with silylsilatrane,7m in which we proposed the reaction proceeds through the four-membered transition state. The screening of bases disclosed Et₃N to be ineffective in the current transformation (entry 7). Lithium acetate or pivalate showed a lower efficiency compared to benzoate (entries 8, 9). A much lower yield was observed with lithium carbonate, indicating the importance of carboxylate bases (entry 10). Sodium and potassium benzoates did not give better yields than lithium benzoate (entries 11, 12). Other solvents were also examined. The reaction efficiency is similar in NMP (entry 13), and a slower reaction was observed in DMF (entry 14). The reaction completely stopped in 1,4-dioxane (entry 15). Under strictly dehydrated conditions, the reduced conversion rate was observed to indicate the importance of water for the

reproducible results in the current transformation, though the exact reason is unclear. Thus, the conditions in entry 1 were confirmed to be optimal.

Table 2 shows the substrate scope under the optimized conditions. Silvlation products were obtained in good yields when the para position was substituted with electronwithdrawing groups such as acetyl (6b¹¹), cyano (6c¹²), benzoyl (6d), CF₃ (6e), and formyl (6f). A naphthyl substituent was also applicable (6g). It is intriguing that the Bpin substituent in 6h survived the reaction conditions although some decomposition, probably at the boron unit, was observed during purification on silica gel. Fluoro-substituted product 6i was obtained albeit with a loss of the material and was obtained in only a 24% yield. With nitro group, none of product 6j was observed while most of the aryl chloride starting material was recovered. With an electrondonating methyl (6d) or methoxy (6h) substituents, conversion was very sluggish. Product bearing *m*-ethoxycarbonyl group **6n** or *m*-cyano group **60** was obtained in moderate yield. Ortho substitution seems to be retarding the silylation and no ethoxycarbonyl product 6p was observed. Heteroaromatic substrate (2-chloroquinoline) or alkenyl chloride (4-tert-butyl-1chlorocyclohex-1-ene) gave no corresponding silylated product.

Other dialkoxydisilanes could also be used for this silylation reaction (Scheme 2). 1,2-Diethoxy-1,1,2,2-tetramethyldisilane **2**, which was reactive under Denmark's conditions^{5d} reacted similarly under our optimized conditions. However, product **3** was not stable enough for chromatographic purification, and the yield (61%) was determined by gas chromatography. Also, we have synthesized 1,2-bis(adamantan-1-yloxy)-1,1,2,2-tetramethyldisilane (**7**) as a crystalline dialkoxydisilane.¹³ Under the standard conditions, **7** could also be applied to the silylation reaction of **4a** to afford **8** in 46% yield.

Bromides and iodides were examined for our silylation reaction (Table 3). Under the standard conditions for aryl chloride, bromide **9** and iodide **10** were converted to the corresponding arylsilane **6a** only in 27% and 3% respective yields (entries 1, 2). In the presence of additional LiCl, yields were slightly improved (27% to 48% for bromide **9** and 3% to 21% for iodide **10**) (entries 3, 4). The reason for the importance of chloride ion is not clear in the current state.



Table 3 Applications to Aryl Bromide and Iodide.



^aDetermined by ¹H NMR analysis using mesitylene as an internal standard.

A plausible reaction mechanism for the silylation is shown in Figure 1. Pd(0) species **11** generated from the precatalyst would be subjected to oxidative addition to yield **12**. Maiti reported a mechanistic investigation by DFT calculations for his palladiumcatalyzed C-H silylation with hexamethyldisilane, in which the chloride substituent on Pd(II) was replaced with a carboxylate to facilitate the activation of the disilane to give a Pd(IV) intermediate.¹⁴ Keeping in mind that carboxylate bases were indispensable for the current silylation, the reaction of **12** with disilane **5** would similarly form Pd(IV) carboxylate species **13** and reductive elimination to generate arylsilane **6**.¹⁵ Finally, reductive elimination from **14** would generate silyl carboxylate **15** and Pd(0) species **11** to close the catalytic cycle.



In conclusion, conditions for palladium-catalyzed silylation reaction between aryl chlorides and bulky dialkoxydisilanes were developed. These NHC-Pd-catalyzed conditions newly allowed the introduction of bulky *tert*-alkoxysilyl groups to arenes bearing electron-neutral or -withdrawing substituents.

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Supporting Information

Supporting information for this article is available online

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- (9) Ethyl 4-(tert-butoxydimethylsilyl)benzoate (6a); Typical procedure An oven-dried 30-mL Schlenk tube was charged with LiOBz (96.0 mg, 0.75 mmol), SingaCycle-A3 (10.0 mg, 0.015 mmol), and DMA (1.5 mL) under nitrogen atmosphere. Ethyl 4-chlorobenzoate (4a, 92.3 mg, 0.50 mmol), 1,2-di-tert-butoxy-1,1,2,2-tetramethydisilane (5, 235 µL, 0.75 mmol), and H₂O stock solution (0.17 M in DMA, 0.15 mL, 25 µmol) were sequentially added to the mixture. DMA (1.35 mL) was added to wash the inner side of the tube. The reaction mixture was stirred at 100 $^{\circ}\mathrm{C}$ for 12 h and quenched with sat. aq. NaHCO3 (10 mL). The mixture was poured into a separatory funnel with AcOEt (20 mL) and partitioned. The organic phase was collected and washed with sat. aq. $\ensuremath{\mathsf{NaHCO}_3}$ (10 mL), sat. aq. NH₄Cl (10 mL), brine (10 mL), dried over anhydrous Na₂SO₄ (ca. 5 g), filtered, and concentrated in vacuo. The residue was purified by column chromatography on silica gel with an eluent (hexane and hexane/AcOEt = 100/1) to afford 6a (105.6 mg, 0.38 mmol, 75%) as colorless oil. ¹H NMR (CDCl₃): δ 8.01 (d, J = 8.2 Hz, 2H), 7.67 (d, J = 8.2 Hz, 2H), 4.38 (q, J = 6.9 Hz, 2H), 1.39 (t, J = 6.9 Hz, 3H), 1.25 (s, 9H), 0.39 (s, 6H); ¹³C NMR (CDCl₃): δ 167.0, 146.8, 133.4, 131.0, 128.6, 73.2, 61.0, 32.2, 14.5, 1.5; HRMS calcd for C15H25O3Si [(M+H)+]: 281.1567, found: 281.1569.
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- (11) 1-(4-(tert-Butoxydimethylsilyl)phenyl)ethan-1-one (6b)
 - Reaction time was 12 h. Column chromatography with an eluent (hexane to hexane/AcOEt = 30/1) afforded **6b** as colorless oil (85.9 mg, 0.34 mmol, 69%) from **4b** (77.3 mg, 0.50 mmol). ¹H NMR (CDCl₃): δ 7.92 (d, *J* = 8.2 Hz, 2H), 7.70 (d, *J* = 8.2 Hz, 2H), 2.61 (s, 3H), 1.26 (s, 9H), 0.39 (s, 6H); ¹³C NMR (CDCl₃): δ 198.6, 147.3, 137.5, 133.7, 127.3, 73.2, 32.2, 26.8, 1.5; HRMS calcd for C₁₄H₂₃O₂Si [(M+H)⁺]: 251.1462, found: 251.1461.
- (12) 4-(tert-Butoxydimethylsilyl)benzonitrile (6c)
 - Reaction time was 12 h. Column chromatography with an eluent (hexane to hexane/AcOEt = 100/1) afforded **6c** as colorless oil (89.6 mg, 0.38 mmol, 77%) from **4c** (68.8 mg, 0.50 mmol). ¹H NMR (CDCl₃): δ 7.68 (d, *J* = 8.2 Hz, 2H), 7.62 (d, *J* = 8.2 Hz, 2H), 1.26 (s, 9H), 0.39 (s, 6H); ¹³C NMR (CDCl₃): δ 147.4, 133.9, 131.1, 119.3, 112.7, 73.4, 32.2, 1.5; HRMS calcd for C₁₃H₂₀NOSi [(M+H)+]: 234.1309, found: 134.1304.
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